

DEC 30 2005

K053303

EXHIBIT 2
510K Summary

Cardicom
1260 Park Road
Chanhassen, MN 55317
Toll-Free: 888-243-8881
Tel: 952-474-4149
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Contact: Daniel Cosentino, President

Date: November 19, 2005

1. **Identification of the device**
Proprietary-Trade Name: Cardicom Commander III Non-Invasive Automated Blood Pressure Monitor
Classification Name: DXN, SYSTEM, MEASUREMENT, BLOOD-PRESSURE, NON-INVASIVE,
Common/Usual Name: Noninvasive Blood Pressure Measurement System
2. **Equivalent legally marketed devices**
This product is similar in function and design to predicate K043096 Cardicom LLC Commander II.
3. **Indications for Use (intended use).** The Commander III device is for use by patients to collect and transmit general health questions and patient vital sign data (such as weight, blood pressure, glucose, pulse oximetry, peak flow) between the patient, typically at home, and a health care professional at a remote site. Contraindications, Precautions and Warnings: The Commander III device makes no interpretation, evaluation, medical judgments or recommendations for treatment. Clinical judgment and experience are required to check and interpret the information transmitted. The Commander III is not intended as a substitute for medical care
4. **Description of the Device.** Commander III is similar to a simple personal computer with a modem that stores and transmits data. Commander III connects to a user's telephone line at home. It has a display that asks health related questions to which the user can respond 'Yes', 'No', or select from a list. It contains an integrated blood pressure meter. It also has inputs for devices such as weight scales, glucometers, peak flow meters and pulse oximeters. These devices download data through a RS232 connection. The functionality of these devices has not been modified; they are used as supplied from the manufacturer.
5. **Safety and Effectiveness, comparison to predicate device**
The results of bench, laboratory, and clinical testing indicates that the new device is as safe and effective as the predicate device.
6. **Comparison Table**

Comparison matrix – new vs. Predicate device Characteristic	K043096 Cardiocom LLC Predicate Device	Submission Device
Operating Principle	Oscillometric automated blood pressure monitoring	SAME
Product Name	Commander II	Commander III
Classification Name	Noninvasive blood pressure measurement system	Noninvasive blood pressure measurement system
Description	The Commander II is an automated device that connects to the user's telephone system at home. It has a display that asks the user health related questions and has inputs for devices such as weight scales, blood pressure meters, and other vital sign measurement devices.	The Commander III is an automated device that connects to the user's telephone system at home. It has a display that asks the user health related questions and has inputs for devices such as weight scales, Glucometers, and other vital sign measurement devices. <u>The Commander III has a built in NIBP meter.</u>
Intended Use	The Commander II device is for use by patients to collect and transmit general health questions and patient vital sign data (such as weight, blood pressure, glucose, pulse oximetry, peak flow) between the patient, typically at home, and a health care professional at a remote site.	The Commander III device is for use by patients to collect and transmit general health questions and patient vital sign data (such as weight, blood pressure, glucose, pulse oximetry, peak flow) between the patient, typically at home, and a health care professional at a remote site.
Blood Pressure Meter	A&D Medical UA-767PC Blood Pressure Monitor for Telemonitoring	BUILT IN Blood Pressure Meter designed by Cardiocom.
Transmission	Telephone line	Telephone line
Intended Users	Home users and health care professionals	Home users and health care professionals
Site of Use	Home: Clinic or remote site	Home: Clinic or remote site
Measurements	<ul style="list-style-type: none"> - Blood pressure - Pulse oximeter - Peak flow meter - Glucose - Weight 	<ul style="list-style-type: none"> - Blood pressure - Pulse oximeter - Peak flow meter - Glucose - Weight
Software	Embedded Firmware and Patient database	embedded Firmware and Patient database
Case	Plastic Wedge shape	Plastic Wedge shape
User interface	Keypad overlay with YES, NO up	Keypad overlay with YES, NO

Comparison matrix – new vs. Predicate device Characteristic	K043096 Cardiocom LLC Predicate Device	Submission Device
	arrow and down arrow	up arrow , down arrow, and selection buttons
Communication	Multiple serial RS-232 ports and telephone modem Part 68 approved	Multiple serial RS-232 ports and telephone modem Part 68 approved
Power supply	External AC to DC supply	External AC to DC supply
Electrical Safety	UL/IEC 60601-1	UL/IEC 60601-1
Electromagnetic compatibility	IEC 60601-1-2	IEC 60601-1-2

7. **Conclusion**

After analyzing bench, test laboratory, and clinical testing data, it is the conclusion of Cardiocom that the Cardiocom Commander III is as safe and effective as the predicate device, has essentially no significant technological differences, and has no new indications for use, thus rendering it substantially equivalent to the predicate device.



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration
9200 Corporate Boulevard
Rockville MD 20850

DEC 30 2005

Cardiocom LLC
c/o Mr. Mark Job
Regulatory Technology Services, LLC
1394 25th Street NW
Buffalo, MN 55313

Re: K053303

Trade Name: Cardiocom Commander III
Regulation Number: 21 CFR 870.1130
Regulation Name: Non-Invasive Blood Pressure Measurement System
Regulatory Class: Class II (two)
Product Code: DXN
Dated: December 12, 2005
Received: December 13, 2005

Dear Mr. Job:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

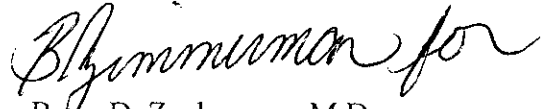
If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

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Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050. This letter will allow you to begin marketing your device as described in your Section 510(k) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Office of Compliance at (240) 276-0120. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 443-6597 or at its Internet address <http://www.fda.gov/cdrh/industry/support/index.html>.

Sincerely yours,

A handwritten signature in black ink, appearing to read "B. Zuckerman for".

Bram D. Zuckerman, M.D.

Director

Division of Cardiovascular Devices

Office of Device Evaluation

Center for Devices and

Radiological Health

Enclosure

Indications for Use

510(k) Number (if known) _____

Device Name: Cardiocom Commander III

Indications For Use:

The Commander III device is for use by patients to collect and transmit general health questions and patient vital sign data (such as weight, blood pressure, glucose, pulse oximetry, peak flow) between the patient, typically at home, and a health care professional at a remote site.

Contraindications, Precautions and Warnings:

The Commander III device makes no interpretation, evaluation, medical judgments or recommendations for treatment. Clinical judgment and experience are required to check and interpret the information transmitted. The Commander III is not intended as a substitute for medical care.

Prescription Use X AND/OR Over-The-Counter Use _____
(Part 21 CFR 801 Subpart D) (21 CFR 807 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of Device Evaluation (ODE)

B. Hammerman
(Division Sign-Off)
Division of Cardiovascular Devices
510(k) Number K053303

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